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CD86 Has Sustained Costimulatory Effects on CD8 T Cells

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CD80 and CD86 both costimulate T cell activation. Their individual effects in vivo are difficult to study as they are coordinately up-regulated on APCs. We have studied mice expressing rat insulin promoter (RIP)-CD80 and RIP-CD86 on the NOD and NOD.scid genetic background to generate in vivo models, using diabetes as a readout for cytotoxic T cell activation. Accelerated spontaneous diabetes onset was observed in NOD-RIP-CD80 mice and the transfer of diabetes from 6-wk-old NOD mice to NOD.scid-RIP-CD80 mice was greater compared with NOD-RIP-CD86 and NOD.scid-RIP-CD86 mice, respectively. However, the secondary in vivo response was maintained if T cells were activated through CD86 costimulation compared with CD80. This was demonstrated by greater ability to cause recurrent diabetes in NOD-RIP-CD86 diabetic mice transplanted with 6-wk-old NOD islets and adoptively transferred diabetes from diabetic NOD-RIP-CD86 mice to NOD.scid mice. In vitro, CD80 costimulation enhanced cytotoxicity, proliferation, and cytokine secretion in activated CD8 T cells compared with CD86 costimulation. We demonstrated increased CTLA-4 and programmed death-1 inhibitory molecule expression following costimulation by both CD80 and CD86 (CD80 > CD86). Furthermore, T cells stimulated by CD80 were more susceptible to inhibition by CD4+CD25+ T cells. Overall, while CD86 does not stimulate an initial response as strongly as CD80, there is greater sustained activity that is seen even in the absence of continued costimulation. These functions have implications for the engineered use of costimulatory molecules in altering immune responses in a therapeutic setting. The Journal of Immunology, 2007, 179: 5936–5946.

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† Abbreviations used in this paper: PD-1, programmed death-1; RIP, rat insulin promoter.
of use when expressed ectopically in tumors for enhancing immune responses. Much less is known of CD86 in this context. The rat insulin promoter (RIP) has been used to direct expression of CD80 or CD86 costimulatory molecules to insulin-producing islet β cells, which do not normally express any costimulatory molecules. In conjunction with endogenous MHC class I molecules, islet β cells are effectively transformed into APCs for CD8 T cells. Model systems using RIP-CD80 and RIP-CD86 have been developed to test whether tissue expression of costimulators on islet β cells would break tolerance and cause diabetes. The experiments showed that, by itself, T cell activation with CD80 costimulation on a B6 or bnl genetic background is not sufficient to cause disease (23–25). However, the presence of other factors, such as the coexpression of a viral protein or MHC class II, IE, could stimulate diabetes (26–28). Coexpression of cytokines TNF-α or IL-2 also precipitated diabetes (23, 28). Furthermore, the expression of CD80 on the NOD genetic background caused accelerated diabetes, as early as 3 wk of age (25, 29, 30). The effect of the MHC in this acceleration is particularly important as CD80 expression on the nonobese-resistant and congenic B6R7 mice also facilitates spontaneous diabetes in these diabetes-resistant strains (31). CD86 has been less studied in these models. When CD86 was expressed on the pancreatic islet β cells on the B6 genetic background, T cell tolerance to β cells was maintained and the mice did not develop diabetes (32). However, there was peri-islet infiltration with IL-4-producing cells in older mice. The effect of CD86 expression on the NOD genetic background was not previously reported.

In this study, we directly compared the effects of CD80 and CD86 costimulation on CD8 T cell activation using diabetes onset as an in vivo read out. We report that CD86 activates CD8 T cells with less potency during primary activation in vivo compared with CD80. In vitro studies, using CD80- or CD86-expressing islets as APCs, demonstrated less enhanced proliferation and cytotoxicity of CD86-activated CD8 T cells compared with CD80. However, CD80 costimulation also leads to greater increases of the inhibitory molecules CTLA-4 and PD-1 which could down-regulate CD8 T cell function. Furthermore, CD8 T cells activated by CD80 are more susceptible to inhibition by CD4+CD25+ T cells. These cells require continued costimulatory activity for maintenance of activation, whereas CD86 costimulation leads to lower but more sustained activity of CD8 T cells.

Materials and Methods

Mice

Two founder lines of C57BL6-RIP-CD86 mice, lines 12 and 46, which expressed human CD86 on pancreatic β cells at medium and high levels, respectively (32), were backcrossed to NOD/caj mice (provided by the late C. Janeway, Yale School of Medicine, New Haven, CT) for at least 14 generations and were designated lines 12B and 46K. NOD-RIP-CD80 mice (29, 30) were backcrossing C57BL6-RIP-CD80 mice to NOD/caj mice for at least three generations. NOD.scid-RIP-CD80 and NOD.scid-RIP-CD86 mice were generated by breeding NOD-RIP-CD80 or NOD-RIP-CD86 mice with NOD.scid mice. Transgene-positive F1 mice were backcrossed to NOD.scid mice and transgene-positive scid homozygous F1 mice were generated. These NOD.scid-RIP-CD80 mice were backcrossed nine generations to NOD.scid mice. NOD.scid-RIP-CD86 mice were derived from RIP-CD86 mice previously backcrossed 14 generations to NOD mice and therefore no further backcross was required.

Insulin-specific TCR-transgenic mice were generated by isolating TCR genomic DNA from G9C8 cloned T cells (33), which have previously been shown to have specific reactivity to amino acids 15–23 of the insulin B chain (34). TCR α (V1e1851 J1e18 Cα) and β (Vβ6S1 Dβ1.1 Jβ2.3 Cα1) chain DNA was purified and cloned into pTeras+ and pTβcas+ constructs, respectively, provided by D. Mathis (Joslin Diabetes Center, Harvard Medical School, Boston, MA) (35). The cloned constructs were injected directly into NOD OVA to generate independent TCR α and β founder lines, which were intercrossed to produce αβ TCR-transgenic mice (G9.NOD, G9Ca−/−). NOD mice were generated by crossing the αβ TCR-transgenic mice to NOD.Ca−/− mice (>20 generations backcross to NOD mice). The mice were specific pathogen free and housed in barrier rooms. All procedures were performed in accordance with U.K. Home Office-approved protocols.

Genotyping of mice for expression of CD80 or CD86 using PCR

Tail-tip DNA from NOD-RIP-CD80 and NOD-RIP-CD86 DNA was analyzed in a standardized PCR. The samples were amplified for 35 cycles of 30 s denaturing at 95°C, 1 min annealing at 55°C, 1 min extension at 72°C and a final extension of 10 min. Primers B7-311 (TGA AGC CAT GGG CCA CAC) and B7-1192 (GAC ACT GTT ATA CAG GAG) produced an 880-bp band for CD80 and primers B70–480 (CCC ACA GGA ATG ATT CGC ATC) and B70–1009 (TCA CTC TCT TCC TCT TCC ATT GTG) produced a 530-bp band for CD86 in transgene-positive mice. Primers were synthesized by Invitrogen Life Technologies.

Detection of CD80 and CD86 from islet cDNA

Pancreata from eight NOD-RIP-CD80 and eight NOD-RIP-CD86 mice at 6 wk of age were harvested and islets were isolated as described. RNA was extracted from the islet samples using TRIzol (Invitrogen Life Technologies). cDNA was generated from the RNA samples according to the manufacturer’s instructions. Concentrations of cDNA were measured using a spectrophotometer and 1/2 dilutions of cDNA (starting at 2 μg) were amplified using primers for CD80 and CD86 as described above. GAPDH was used as a control (GAPDH forward: GGT CAT CAT CTC CGC CCC TTC and the AP substrate kit, VectorRed (Vector Laboratories). VectorRed working solution, made up in 100 mM Tris-HCl (pH 8.4) with the addition of levamisole (Vector Laboratories), was used in accordance with the manufacturer’s instructions. Sections were incubated in the dark for a preoptimized time and counterstained with hematoxylin (Vector Laboratories) before mounting under coverslips with Hydromount (National Diagnostics). Insulitis was measured in diabetic, 6- and 12-wk-old NOD, NOD-RIP-CD80, and NOD-RIP-CD86 mice. Sections were stained with H&E (Vector Laboratories) as described (29, 30) and 3–200 islets from four mice per group were scored for insulitis. Score 0, no infiltration; 1, peri-insulitis; 2, ≤50% infiltration; 3, 50–90% infiltration; and 4, ≥90% infiltration.

Diagnosis of diabetes

Groups of NOD-RIP-CD86 lines, NOD-RIP-CD80, and female NOD-RIP-CD86 transgene-negative mice were monitored weekly for spontaneous development of diabetes over a period of 25 wk. Diabetes was detected by initially testing for glycosuria (Bayer Diastix). Diagnosis was confirmed by blood glucose measurements >14 mM/L. Mice from adoptive transfer and transplant experiments were tested two to three times weekly for 112 days or until diabetic.

Adoptive transfer of diabetes

A total of 20 × 106 splenocytes at 106 cells/ml, isolated from diabetic NOD, NOD-RIP-CD80, and NOD-RIP-CD86 mice or 6-wk-old NOD mice or 104 purified CD8 T cells from G9Ca−/−.NOD mice were i.v. injected into 6- to 8-wk-old NOD.scid, NOD.scid-RIP-CD80, or NOD.scid-RIP-CD86-transgenic mice.

Murine islet isolation and transplantation

Perfusion of murine pancreas followed an adapted version based on a modification of the method established by Gotoh et al. (36). In brief, pancreata from NOD.scid, NOD.scid-RIP-CD80, NOD.scid-RIP-CD86, and diabetic transgenic mice were perfused with 2–3 ml of collagenase P (1 mg/ml; Boehringer Mannheim). Bile ducts were clamped off at the duodenal insertion to allow cannulation and specific perfusion to the pancreas. Perfused pancreata were digested for a batch-specific optimized time and collagenase activity was stopped with ice-cold HBSS, 10 mM HEPES, 5% FCS (Invitrogen Life Technologies). Digested pancreata were passed through a 100-μm strainer and centrifuged through layers of 25, 23, 20,
and 11% Ficoll in HBSS-10 mM HEPES. Islets were collected at the interface between layers and for in vitro assays were handpicked under a dissecting microscope into RPMI complete medium (RPMI 1640–5% FCS, 50 μM 2-ME, 2 mM l-glutamine, 100 U/ml penicillin, 100 μg/ml streptomycin, 0.25 μg/ml amphotericin; Invitrogen Life Technologies). When used for culture, the islets were irradiated with 4200 rad. For extraction of islet-infiltrating cells, the islets were trypsinized and filtered through a 70-μm cell strainer before flow cytometric analysis.

**Flow cytometry**

Single-cell suspensions of 0.1–1 × 10^6 cells were stained for various cell surface markers (rat anti-murine CD8-allophycocyanin; CD25, CD62L, CD95L, 4-1BB, ICOS, and PD-1-PE; CD107a and CD95-FITC). mAbs were blocked with anti-CD16 (2.4G2, provided by the late C. A. Janeway) and cells were stained for surface markers and costimulatory molecules. Intracellular staining was performed for CTLA-4 after cell surface staining following BD Pharmingen’s protocol. Intracellular staining for FoxP3 was performed using a FoxP3 staining kit (eBioscience/Insight Technology). The samples were acquired by flow cytometry using a FACSCalibur, results were analyzed using FlowJo software. All Abs had been previously titrated to determine optimal concentration and all were purchased from BD Pharmingen, except anti-murine CD8-allophycocyanin (Caltag Laboratories).

**Analysis of cytokine secretion in T cell cultures**

Supernatants from cell stimulation assays were harvested and analyzed for the presence of secreted cytokines using a multiplex fluorescent bead immunoassay from Bender Medsystems (Caltag Laboratories) (mouse Th1/Th2 cytokines: IFN-γ, IL-10, IL-2, IL-4, IL-5, IL-6, IL-10, IL-17, TNF-α, and GM-CSF). Measurements were performed as described in the manufacturer’s instructions. Data were analyzed using BMS FlowCytomix Software, which was based in WinMDI and Microsoft Excel.

**Proliferation, cytokine secretion, and cytotoxicity assays**

Insulin-reactive CD8 T cells (3 × 10^5) from 5- to 8-wk-old G9C+/−NOD mice were cocultured with 25 irradiated islets from NOD.scid, NOD.scid-RIP-CD80, or NOD.scid-RIP-CD86 mice in RPMI complete medium. CD8 T cells were purified from G9C+/−NOD splenocytes using positive selection beads (Miltenyi Biotech) with >95% purities. Islets and CD8 T cells alone were used as controls. After 2 days of incubation, supernatants were taken for cytokine secretion analysis and cells were either pulsed with [3H]thymidine for 14 h to test proliferation or were used in cytotoxicity assays. Stimulated CD8 T cells were harvested and further cocultured with 10^5 51Cr-sodium chromate (Amersham) labeled P815 cells together with insulin peptide (B15–23) at an E:T ratio of 10:1 for 16 h. Specific lysis was calculated as (cytotoxic release – min)/(max – min)) × 100%, where the minimal release (min) corresponds to the spontaneous lysis, and the maximal lysis corresponds to lysis induced by addition of hydrochloric acid (max).

**CD4⁺ CD25⁺ T cell depletion**

CD25⁺ T cells were depleted using the CD25 depletion kit (Miltenyi Biotech) according to the manufacturer’s instructions. The depletion removed representative of three individual mice per strain. B, Comparative expression of CD80 and CD86 levels was tested using PCR amplification of 2-fold dilutions of cDNA (0.5–9.8 × 10^−4 μg) extracted from islets isolated from eight mice per strain. GAPDH was used as a housekeeping control. C, Densitometry confirmed similar expression of CD80 and CD86 expression. D, Male and female NOD-RIP-CD80, NOD-RIP-CD86 (12B and 46K), and transgene-negative mice were monitored weekly for diabetes onset by testing for glycosuria and confirmed by testing blood glucose levels (>14 mM/L). Log-rank tests showed no statistically significant difference between 12B and 46K mice (p = 0.28) but significant differences were found between NOD-RIP-CD80 and NOD-RIP-CD86 (12B and 46K) mice (p = 0.002). E, Islet infiltrates from diabetic NOD-RIP-CD80, NOD-RIP-CD86, and transgene (Tg) negative mice were tested for the presence of CD8, CD4, and CD25⁺ T cells by immunohistochemistry on frozen sections. Sections shown are representative of three individual mice per strain.
diabetes from 7 to 8 wk of age, 50% diabetes incidence between 13 and 46K-transgenic mice showed comparable results with an onset of male diabetes incidence curves of NOD-RIP-CD86 12B and CD86-transgenic mice (data not shown). Combined male and female (12B) and high (46K) expressing founder lines of NOD-RIP-CD80 mice had levels of infiltration compared to 90 and 50% in NOD-RIP-CD86 and NOD-RIP-CD80 islets, respectively. Values of $p \leq 0.0009$ for all three comparisons. The difference in NOD-RIP-CD80 and NOD-RIP-CD86 islet infiltration compared to transgene negative mice was statistically different by Fisher’s exact test ($p = 0.0002$ and 0.0012, respectively). Similar degrees of infiltration between NOD-RIP-CD80 and NOD-RIP-CD86 mice at 12 wk corresponded with the convergence of diabetes incidence in these mice at 12 wk of age. In diabetic mice, there was no difference in degree of insulitis between all strains, where >97% of all islets had high levels of infiltration ($p > 0.3$ for all comparisons).

**Statistical analysis**

Statistical analysis was performed as stated using Graphpad Prism software. Analyses of Kaplan-Meier survival curves used a log-rank test similar to a Mantel-Haenszel test. Fisher’s exact tests were performed for insulin comparisons. For both tests, Bonferroni corrections were made for multiple comparisons.

**Results**

Transgene expression is similar in NOD-RIP-CD86 and NOD-RIP-CD80 mice

A comparative study of spontaneous diabetes incidence was performed using NOD-RIP-CD80 and two founder lines of NOD-RIP-CD86-transgenic mice. Ab staining of pancreatic sections for the presence of transgenes was specific in the appropriate mice and demonstrated similar levels of staining in both NOD-RIP-CD80 and NOD-RIP-CD86 mice (Fig. 1A). A 2-fold limiting dilution PCR of NOD-RIP-CD80 and NOD-RIP-CD86 cDNA demonstrated comparatively similar levels of expression for the 880- and 530-bp products, respectively, related to the control products of the housekeeping gene GAPDH (Fig. 1, B and C).

**NOD-RIP-CD86 mice develop accelerated diabetes but less rapidly than NOD-RIP-CD80 mice**

A previous report demonstrated no difference in the onset of autoimmune diabetes in two founder lines of NOD-RIP-CD86-transgenic mice with differing levels of CD8 expression or a difference between genders (29). Similar to NOD-RIP-CD80 mice, no gender difference in the onset of diabetes was observed between the medium (12B) and high (46K) expressing founder lines of NOD-RIP-CD86-transgenic mice (data not shown). Combined male and female diabetes incidence curves of NOD-RIP-CD86 12B and 46K-transgenic mice showed comparable results with an onset of diabetes from 7 to 8 wk of age, 50% diabetes incidence between 13 and 14 wk and >80% diabetes incidence by 20 wk of age. There was no statistically significant difference in the diabetes incidence between the two lines of NOD-RIP-CD86 (Fig. 1D). In contrast, NOD-RIP-CD80 mice developed diabetes from an earlier age of 4 wk, 50% were diabetic by 13 wk and at 20 wk of age diabetes incidence was 58% (Fig. 1D), which was similar to previously published work (29, 30). In addition, at the point of diabetes onset in NOD-RIP-CD86-transgenic mice, 15% of NOD-RIP-CD80-transgenic mice had become diabetic (Fig. 1D). Log-rank tests comparing NOD-RIP-CD80 and combined 46K and 12B NOD-RIP-CD86 incidence curves showed that diabetes incidence was significantly different ($p = 0.002$). Both NOD-RIP-CD80 and NOD-RIP-CD86 mice had accelerated onset of diabetes compared with 13-wk-old transgene-negative mice.

Expression of RIP-CD80 and RIP-CD86 and acceleration of autoimmune diabetes was associated with higher levels of islet infiltration (Table I) and an increased ratio of CD8 T cells in islets from the islets (Fig. 1, E and F). At 6 wk of age, islet infiltration was most pronounced in NOD-RIP-CD80 mice, where ~50% of islets had high levels of infiltration (scores 2–4), compared with 10% in NOD-RIP-CD86 mice and only low levels (scores 0–1) of infiltration were observed in transgene-negative mice (Table I). By 12 wk of age, the degree of insulitis in NOD-RIP-CD80 and NOD-RIP-CD86 islets was comparable but more insulitis was observed compared with transgene-negative mice. Islet infiltration was similar in all strains after diabetes development (Table I). Islet infiltrates in both diabetic NOD-RIP-CD80 and NOD-RIP-CD86 mice comprised a high proportion of CD8 T cells compared with transgene-negative mice, while the proportion of CD4 T cells and B cells was comparable (Fig. 1E). This was confirmed by flow cytometry showing that isolated islet infiltrates from diabetic mice demonstrated a 3-fold increase in the proportion of CD8 T cells from NOD-RIP-CD80 and NOD-RIP-CD86 mice compared with transgene-negative mice (Fig. 1F).

**Recurrent diabetes occurs at a faster rate in diabetic NOD-RIP-CD86 mice compared with diabetic NOD-RIP-CD80 mice transplanted with healthy NOD islets**

We tested secondary responses of cells from diabetic NOD, NOD-RIP-CD80, and NOD-RIP-CD80 mice in islet transplant experiments. Islets from 6-wk-old NOD mice were transplanted into diabetic NOD, NOD-RIP-CD80, and NOD-RIP-CD80 recipient mice under the kidney capsule. The recipients of the transplants became euglycemic by 48 h. Infiltration and destruction of the transplanted islets recurred with a reversed pattern of onset compared with the natural history experiments (Fig. 2A). Diabetes recurred in NOD mice between 7 and 12 days post-islet transplantation and in NOD-RIP-CD86 mice between 17 and 18 days post-islet transplantation. In contrast, diabetes recurred in NOD-RIP-CD80 mice from 25 days posttransplantation and one recipient even survived for 45 days posttransplantation (Fig. 2A). This suggested that the cells had been “tuned” such that cells from RIP-CD80 mice were less effective than those from RIP-CD86 mice, which in turn were less effective than cells from the NOD mice. Curiously, when transplanted islets were examined for the presence of CD8, CD4, and B cells, all the CD4+CD25 high cells and <5% of the low-staining CD4+CD25+ cells remained after the procedure.
a marked reduction in the CD8:CD4 T cell ratio was observed in NOD-RIP-CD80-transplanted mice (Fig. 2B). The relative number of infiltrating CD4 T cells was at least double compared with infiltrating CD8 T cells in the NOD-RIP-CD80-transplanted mice. In contrast, both NOD and NOD-RIP-CD86 diabetic mice had similar proportions of infiltrating CD8 and CD4 T cells although the NOD-RIP-CD86 islets had a greater total infiltrate. B cells followed a similar pattern of infiltration into islet transplants of diabetic NOD and NOD-RIP-CD86 mice. CD8 T cells from diabetic NOD-RIP-CD86 mice have a greater ability to transfer diabetes compared with CD8 T cells from diabetic NOD-RIP-CD80 mice

Previous studies have shown that splenocytes from diabetic NOD-RIP-CD80 mice have a reduced capacity to transfer diabetes to NOD.scid recipients (F. S. Wong, unpublished data and Ref. 25). In this study, we compared the ability of splenocytes from diabetic NOD-RIP-CD86 and NOD-RIP-CD80 mice to transfer diabetes to NOD.scid recipients. Eighty-nine percent of recipient mice developed diabetes after transfer of NOD-RIP-CD80 diabetic splenocytes. Peri-insulitis consisting of CD8 T cells, CD4 T cells, and B cells was observed in the islets of nondiabetic recipient NOD.scid mice at 112 days posttransfer with NOD-RIP-CD80 or NOD-RIP-CD86 diabetic splenocytes (Fig. 3B). Clearly, transferred splenocytes from diabetic NOD-RIP-CD80 mice were capable of reaching the islets and when
further costimulation was provided by transfer into NOD.scid-RIP-CD80 recipients, these cells were capable of adaptively transferring diabetes in 100% of recipients (Fig. 3C). Thus, cells costimulated by CD80 were CD80 dependent whereas those stimulated by CD86 were not.

**T cell development does not play a role in the difference in T cell activation by CD86 or CD80**

To test that the different outcomes following adoptive transfer were not related to differences in T cell development in the thymus, we used adoptive transfer of splenocytes from 6-wk-old NOD mice into NOD.scid-RIP-CD86 and NOD.scid-RIP-CD80 mice and tested for their ability to develop diabetes (Fig. 4A). A similar pattern of development of diabetes compared with the natural history of diabetes in NOD-RIP-CD80 and NOD-RIP-CD80 mice was observed. One hundred percent of NOD.scid-RIP-CD80 recipients had developed diabetes by 40 days posttransfer, while the earliest time of diabetes onset occurred 45 and 54 days in NOD.scid-RIP-CD80 and NOD.scid mice, respectively. By 70 days after transfer, 90% of NOD.scid-RIP-CD86 recipients had developed diabetes whereas only 45% of NOD.scid recipients had developed disease at this point (Fig. 4A). Thus, this experiment removed the possibility that any differences seen in diabetes onset in the NOD-RIP-CD80 or NOD-RIP-CD80 mice related to expression of the transgenes in the thymus and hence to development of diabetogenic T cells.

Additionally, greater expansion (or recruitment) of CD8 T cells in islet infiltrates was observed in NOD.scid-RIP-CD80 recipients transferred with spleen cells from 6-wk-old NOD mice (Fig. 4B). Islet infiltration was similar at 16 days posttransfer, where ~5–6% CD8 and 10–11% CD4 T cells were observed (Fig. 4B). By 28 days, the proportion of CD8 T cells in NOD-scid-RIP-CD80 recipients had risen to >30%, double that compared with NOD.scid-RIP-CD86 recipients. Conversely, the level of CD4 T cells was increased in the NOD.scid-RIP-CD86 recipient mice (Fig. 4B). Early diabetes onset, increased islet infiltration, and greater numbers of CD8 T cells all suggest that islet expression of CD80 and CD86 enables specific activation of CD8 T cells. The fact that these observations occur at an earlier time point in NOD-RIP-CD80 mice strongly indicates that activation of CD8 T cells through CD80 generates enhanced responses, compared with CD86.

To demonstrate that the expression of CD80 or CD86 could differentially stimulate CD8 T cells directly in our model system, we adaptively transferred purified insulin-specific CD8 T cells directly ex vivo from the splenocytes of G9Ca−/−, NOD-transgenic mice into NOD.scid-RIP-CD80 and NOD.scid-RIP-CD86 recipients. We showed that the naive CD8 insulin-reactive T cells transfer diabetes more rapidly and with a higher final incidence to NOD.scid-RIP-CD80 recipients compared with NOD.scid-RIP-CD86 recipients shown in Fig. 4C.

**In vitro stimulation of lymphocytes with islets from NOD.scid-RIP-CD80 mice induces greater proliferation and cytokine production than islets from NOD.scid-RIP-CD86 mice and nontransgenic mice**

To test the islet reactivity of T cells of known specificity, CD8 T cells were isolated from insulin-specific CD8 TCR (G9.Ca−/−) transgenic mice and in vitro assays were performed using isolated islets as both a source of CD8 T cell-specific APCs as well as insulin peptides. Proliferation assays clearly demonstrated that stimulation of insulin-specific CD8 T cells using islets from NOD.scid-RIP-CD80 mice generated heightened activity, ~4-fold.

**FIGURE 4.** Accelerated diabetes in RIP-CD80 and RIP-CD86 mice is not due to differences in thymic T cell development. A. Adoptive transfer of diabetes into NOD.scid, NOD.scid-RIP-CD80, and NOD.scid-RIP-CD86 mice showed a similar pattern of diabetes development to the natural history of diabetes in NOD, NOD-RIP-CD80, and NOD-RIP-CD86 mice. A total of 20 × 10⁶ splenocytes from 6-wk-old mice were transferred into NOD.scid, NOD.scid-RIP-CD80 (NSCD80), and NOD.scid-RIP-CD86 (NSCD86) mice and monitored weekly for diabetes development. The difference in the time taken to develop diabetes and the final incidence of diabetes was statistically significant (p < 0.0001). B. Islet expression of CD80 promotes increased islet infiltration of CD8 T cells in NOD.scid-RIP-CD80 recipients over time. Islet-infiltrating cells isolated from NOD.scid-RIP-CD80 and NOD.scid-RIP-CD86 islets after transfer with splenocytes from 6-wk-old mice were stained for CD8 and CD4 T cells. Top panels, Dot plots for cells from 16 days posttransfer; bottom panels, 28 days posttransfer. Islets from three mice per group were used. C. Purified insulin-specific CD8 T cells cause increased incidence of diabetes in NOD.scid-RIP-CD80 recipients compared with NOD.scid-RIP-CD86 recipients. Purified insulin-specific CD8 T cells (10⁵ (>95% pure)) were transferred into NOD.scid-RIP-CD80 (NSCD80) recipients compared with NOD.scid-RIP-CD86 (NSCD86) recipients. The difference in the time taken to develop diabetes and the final incidence of diabetes was statistically significant (p < 0.0001). No diabetes is seen when these cells are transferred to NOD.scid recipients (n = 10, data not shown).
greater compared with proliferation to NOD.scid islets (Fig. 5A). NOD.scid-RIP-CD86 islets also demonstrated enhanced stimulation, albeit to a lesser extent, where proliferation was ~3-fold greater than the proliferation to NOD.scid islets (Fig. 5A). Additionally, it was observed that stimulation of insulin-reactive CD8 T cells with CD80 expressing islets produced increasing levels of IL-2 and IFN-γ secretion over time, while CD86 stimulation, under these conditions, generated very low levels of IL-2 and negligible amounts of IFN-γ (Fig. 5B). In addition, the production of cytokines IL-1α, IL-4, IL-5, IL-6, IL-10, IL-17, TNF-α, and GM-CSF was also tested but these were not detected (data not shown). Higher levels of CD25 and reduced expression of CD62L on CD8 T cells activated by NOD.scid-RIP-CD80 islets compared with NOD.scid-RIP-CD86 islets reflected enhanced T cell activation through CD80 (Fig. 5C).

**FIGURE 5.** In vitro activation of insulin-reactive CD8 T cells by CD80 results in enhanced proliferation, cytokine production, and change in expression of activation markers compared with CD86 stimulation. A, Naive insulin-reactive CD8 T cells proliferated to islets from all strains, but CD80 costimulation induced the highest level of proliferation in a 3-day culture, followed by CD86 stimulation. B, Levels of IL-2 and IFN-γ secretion were measured in supernatants taken from proliferation assays of insulin-reactive CD8 T cells cultured with NOD.scid-RIP-CD80 and NOD.scid-RIP-CD86 islets. C, Cell surface expression of activation markers CD25 and CD62L were measured 2 days after insulin-reactive CD8 T cells were activated by both NOD.scid-RIP-CD80 and NOD.scid-RIP-CD86 islets. Filled histograms represent naive ex vivo insulin-reactive CD8 T cells. Results are representative of three similar assays.

**FIGURE 6.** Stimulation of insulin-reactive CD8 T cells by CD80 produces cells with greater cytotoxic effects compared with stimulation through CD86. A, Cytotoxicity of insulin-reactive CD8 T cells prestimulated with NOD.scid (NS), NOD.scid-RIP-CD80, or NOD.scid-RIP-CD86 islets was measured as specific lysis of insulin B15–23-pulsed 51Cr-labeled P815 cells. B, Levels of cell surface CD107a was measured on prestimulated insulin-reactive CD8 T cells in response to P815 cells in the presence or absence of insulin B15–23. C, Expression levels of death receptor molecules CD95 and CD95L were monitored after activation of insulin-reactive CD8 T cells with islets from NOD.scid-RIP-CD80 or NOD.scid-RIP-CD86 mice. Results are representative of two similar assays.

In vitro stimulation of lymphocytes with islets from NOD.scid-RIP-CD80 mice alters the threshold of activation of cytotoxic responses compared with islets from NOD.scid-RIP-CD86 mice and nontransgenic mice

The cytotoxicity of insulin-reactive CD8 T cells after activation with islets from the different transgenic strains showed that CD80 costimulation induced greater levels of effector function with specific target cell lysis reaching a maximum of 68%, compared with 34 and 31% for CD86 and NOD.scid stimulated cells, respectively (Fig. 6A). Cells stimulated by both CD80 and CD86 respond to lower peptide concentrations, but to a much greater degree in CD80-costimulated CD8 T cells (Fig. 6A), indicating lowering of activation threshold.

With insulin peptide at $3.9 \times 10^{-3} \mu g/ml$, CD8 T cells prestimulated with CD80 islets caused ~34% specific lysis, at which point specific lysis was only 12 and 4% by CD86 and NOD.scid prestimulated CD8 T cells, respectively. For CD86 prestimulated CD8 T cells to cause the same level of cytotoxic killing, 0.5 µg/ml peptide was required (Fig. 6A). Thus, >100 times more peptide was required for CD86-stimulated cells to cause the same level of cytotoxicity as CD80-stimulated cells.

Furthermore, the results demonstrate that specific cell lysis could be perforin/granzyme or death receptor mediated as increased levels of CD107a were found on CD8 T cells following coculture with insulin peptide-coated P815 cells (Fig. 6B) as well as elevated expression of CD95 on islet-activated CD8 T cells, although the overall level of CD95L was low (Fig. 6C).
We found that after depletion of the CD4 \(^+\) T cells, spleenocytes from diabetic NOD.scid-RIP-CD80 were able to transfer diabetes at a similar rate to cells obtained from diabetic NOD.scid-RIP-CD86 mice (Fig. 8B).

**Discussion**

Costimulation is essential for the full activation of CD8 T cells. In this study, we have aimed first to study the effect of CD80 (B7.1) and CD86 (B7.2) individually, a feature not possible in natural APCs as both are coexpressed and regulation is interdependent. Second, we are studying cells that respond to an important autoantigen, insulin, that is presented endogenously in the islets. Third, the strength of signal 1 is also an important factor. Using this model, we are able to use the ability of the islet to naturally present the range of Ag concentrations that the cells would encounter in vivo. The ability of CD8 T cells to cause diabetes is a complex process and our model allows us to test effects on cell expansion, cytokine production, cytotoxicity in vitro and in vivo as well as homing in vivo. We are using diabetes as an in vivo readout to study the individual effects of costimulatory molecules and not the development of natural diabetes, as islet \(\beta\) cells do not normally express costimulatory molecules (37).
In this study, we have demonstrated a clear difference in the activation of CD8 T cells after CD80 or CD86 costimulation. In vivo, we observed accelerated diabetes in RIP-CD80 mice (4 wk) as noted in previous studies (25, 29, 30) as well as in RIP-CD86 mice (7 wk) compared with transgene-negative mice (13 wk). The greater acceleration of diabetes incidence in RIP-CD80 mice was not likely to be due to a difference in expression levels of CD80 compared with CD86. Diabetes development in the medium (12B) and high (46K) expressing NOD-RIP-CD86 mice was almost identical, while previous studies showed a similar result with different expression levels in RIP-CD80 mice (29). Additionally, Ab staining of pancreatic sections from NOD-RIP-CD80 and NOD-RIP-CD86 (12B) had similar protein levels, and PCR products from cDNA isolated from islets of these transgenic mice demonstrated comparable levels of expression. Hence, we infer that accelerated diabetes was due to enhanced stimulation of T cells by CD80 compared with CD86.

The proportion of CD8 T cells in islet infiltrates was increased in diabetic NOD-RIP-CD80 and NOD-RIP-CD86 mice compared with diabetic transgene-negative littermates. However, proportions of B and CD4 T cells were similar in all the diabetic mice examined, which suggested that costimulation and MHC class I on islet β cells specifically activated CD8 T cells. We further demonstrated that CD80 enhanced the proportion of CD8 T cells in islet infiltrates more than CD86 costimulation. NOD.scid-RIP-CD80 and NOD.scid-RIP-CD86 mice i.v. transferred with splenocytes from 6-wk-old NOD mice showed similar proportions of CD8 and CD4 T cells in islet infiltrates after 16 days but after 28 days, twice the CD8 T cells were present in NOD.scid-RIP-CD80 islets compared with NOD.scid-RIP-CD86. This suggested that CD8 T cells were being further activated in situ and hence proliferating in the islets themselves.

Further evidence that the expression of CD80 and CD86 expressed on the islets directly stimulate CD8 T cells was shown by experiments using purified, naive low-avidity CD8 T cells recognizing insulin peptide B15–23. These cells are unable to transfer diabetes to NOD.scid mice unless they have been preactivated (L. K. Siew, F. S. Wong, unpublished observations). However, they cause diabetes in NOD.scid-RIP-CD80 mice with faster kinetics than NOD.scid-RIP-CD86 mice. No CD4 T cells were transferred in these experiments indicating that the expression of the costimulatory molecules on the islets has a direct effect on CD8 T cells.

When the cells from the diabetic NOD-RIP-CD80 and NOD-RIP-CD86-transgenic mice were subjected to a secondary stimulation, CD86-stimulated cells were more effective in causing recurrent diabetes in both NOD islet-transplanted mice and in NOD.scid recipients receiving cells from diabetic NOD-RIP-CD86-transgenic mice. Although the cells from diabetic NOD-RIP-CD80-transgenic mice caused delayed diabetes in NOD.scid recipients or did not transfer diabetes, when the cells from diabetic NOD-RIP-CD80-transgenic mice were retransferred into NOD.scid-RIP-CD80-transgenic mice, diabetes occurred rapidly and in all mice. This indicated that the threshold of activation of the CD80-stimulated cells was altered such that they could respond in the presence of CD80 but only poorly in the absence of the costimulatory molecule.

To examine the effect of the stimulation by islet Ag with either CD80 or 86, we performed in vitro studies with T cells of a defined specificity that are able to react to the islet Ag, insulin. Islets isolated from both NOD.scid-RIP-CD86 and NOD.scid-RIP-CD80 mice induced higher levels of proliferation in naive insulin-reactive CD8 T cells compared with NOD.scid islets but greater proliferation was seen with RIP-CD80 islets. CD8 T cells secreted limited IL-2 and IFN-γ upon stimulation with RIP-CD86 islets, while RIP-CD80 islets induced production of high levels of both cytokines. Other cytokines were not detected and therefore, we infer that the results seen were not due to production of some protective cytokines. The insulin-reactive CD8 T cells stimulated in vitro with RIP-CD80 islets also demonstrated higher levels of cytoxicity, compared with CD86 stimulation, which in turn was higher than cells stimulated with NOD.scid islets. Additionally, responses were more sensitive in CD8 T cells stimulated with CD80 where ~50% specific lysis occurred with peptide concentrations at 0.01 μg/ml. In contrast, ~20 and 10% lysis was observed in CD86-stimulated and unstimulated CD8 T cells, respectively. The studies also demonstrated increased levels of CD107a and CD95 on the surface of activated insulin-reactive CD8 T cells after stimulation with all sets of islets, suggesting that the cytotoxic effects could be manifested through both the death receptor and perforin/granzyme mechanisms. Even though the levels of CD107a were similar throughout, the cytotoxicity shown by CD80-stimulated cells toward peptide coated targets was much higher. It is conceivable that an increased quantity of perforin and the various granzyme molecules are responsible for activating the caspase pathway within the cytotoxic vesicles of CD80-stimulated cells, although we have no direct evidence for this.

We have clearly shown here that stimulation of naive CD8 T cells in vivo and in vitro by both CD80 and CD86 results in activated cells with a greater potential to proliferate and kill through cytoxic mechanisms. Previously it has been demonstrated that cells from diabetic NOD-RIP-CD80 mice have an impaired ability to transfer diabetes to NOD.scid mice (Ref. 25 and F. S. Wong, unpublished observations) whereas splenocytes from diabetic non-transgenic NOD mice transfer diabetes to 100% of NOD.scid recipients. We confirm these results in respect of cells derived from diabetic NOD-RIP-CD80 mice. In addition, however, we show here although the primary stimulation in the presence of CD86 is not as great as with CD80, the CD86-stimulated cells retain the ability to kill targets and cause diabetes in the absence of continued costimulation, and nearly 90% of recipients become diabetic following adoptive transfer of splenocytes from diabetic NOD-RIP-CD86 mice. Thus, although activation through CD80 results in enhanced proliferation and cytotoxicity in a primary response, the secondary responses are markedly reduced. This was further demonstrated in diabetic NOD, NOD-RIP-CD80, and NOD-RIP-CD86 mice transplanted with islets from 6-wk-old mice, in which grafts transplanted into diabetic NOD-RIP-CD86 mice were destroyed faster compared with the longest survival in NOD-RIP-CD80 mice.

The reduced efficacy of CD8 T cells previously stimulated with CD80 compared with CD86 makes CD86 overall a more effective costimulator. It is likely that more than one mechanism is involved in the reduced secondary stimulation of CD8 T cells by CD80 in this vivo model. These could include:

1) Cells previously stimulated by CD80 are not able to migrate to the relevant site (healthy islets), although this is unlikely because CD8, CD4, and B cells were all present in peri-islet infiltrates in non-diabetic transfer recipients at 10 wk. Similarly, in the transplant experiments, lymphocytes were present as infiltrates in the islet grafts. However, in the transplants, the proportion of CD8 T cells was greatly reduced in NOD-RIP-CD80 recipients. The fact that the cells were able to migrate to the islets but not penetrate and destroy the islet cells suggests that they were not sufficiently activated by the islet cells that did not express the costimulatory molecules.

2) Stronger stimulation of CD8 T cells by CD80 induces islet reactive cells to die by activation-induced cell death. However,
after in vitro stimulation with islets expressing CD80 or CD86, survival of the cells is similar as shown by similar cell numbers and annexin V staining (data not shown). Furthermore, following in vitro stimulation with islets and then a short maintenance period over 3 days in IL-2, a greater proportion of CD80-stimulated CD8 T cells remained viable (data not shown).

3) A possible explanation behind the reduced efficacy of CD80-stimulated CD8 T cells is increased and sustained expression of inhibitory coreceptors, such as CTLA-4 and PD-1, although CD86 also up-regulated these inhibitory receptors. It is well known that CTLA-4 is up-regulated on activation and interacts with CD80 and cells are inhibited as a result of this interaction. However, PD-1 up-regulation has not previously been linked to cells activated by CD80. In this study, we have demonstrated that in vivo, PD-1 is up-regulated on the cell surface of CD8 T cells stimulated by CD80 and CD86 but higher expression is evident at an earlier time in CD80-stimulated CD8 T cells. Additionally, intracellular expression of CTLA-4 is apparent in insulin-reactive CD8 T cells at a greater level after activation through CD80 stimulation compared with CD86 stimulation. We propose that after the initial aggressive activation of CD8 T cells by CD80, the T cells up-regulate inhibitory coreceptors to a greater degree and potentially for a longer period of time, compared with CD86 stimulation. This would enable T cells to be “turned off” and, depending on the duration of inhibitory coreceptor expression, prevent or reduce the possibility of a secondary stimulation. This would most likely be the case when peptides of interest are displayed at endogenous concentrations, such as in the studies presented here. However, by providing increased levels of costimulation, this inhibition can be overcome. We performed in vitro blocking experiments using CD8 cells stimulated by RIP-CD80 and RIP-CD86 islets with Abs to block CTLA-4 or PD-1 and tested in vitro cytotoxicity measuring CD107a but did not observe consistent alteration in the in vitro responses (data not shown). Clearly, it would be interesting to monitor the kinetics of inhibitory receptor expression before, during, and after primary and secondary activation of CD8 T cells through CD80 or CD86 stimulation in vivo. Most recently, it has been shown that in addition to binding to the known inhibitory ligand, CTLA-4 on T cells, B7-1 is also able to directly bind to PD-L1 and deliver an inhibitory signal (38). This interaction could also explain the inhibition observed after the initial activation. It would certainly be of importance also to test for the up-regulation of PD-L1 and the effect of blocking this interaction in this system.

4) A further possibility for the reduced ability of CD80-stimulated cells to cause diabetes in secondary adoptive transfer could be an increase in regulatory T cell inhibition of cells stimulated with CD80 compared with CD86. We measured the percentage of CD4+CD25+ cells in the splenocytes of mice that had become diabetic after the primary transfer of 6-wk-old NOD splenocytes into NOD.scid-RIP-CD86 and NOD.scid-RIP-CD80 mice. The percentage of these cells in the splenocytes was not different between the two types of mice. However, following depletion of the CD4+CD25+ T cells, the cells from the diabetic NOD.scid-RIP-CD80 mice were able to transfer diabetes with similar kinetics to cells from diabetic NOD.scid-RIP-CD86 mice. Thus, this suggests that either an increase in the levels of CD4+CD25+ T cells in the diabetic RIP-CD80 splenocytes were inhibiting the CD80-stimulated CD8 T cells (our evidence did not support this), or the CD8 cells were more susceptible to inhibition by these regulatory cells. Removing the CD4+CD25+ T cells facilitated the secondary adoptive transfer of diabetes by the CD80-stimulated splenocytes. Finally, the use of ectopic expression of costimulatory molecules to boost immune responses is a strategy that has been used for modulating immune responses to tumors. The fact that CD80 stimulates stronger activating responses also has an impact in vivo. The increased stimulation of cells with more rapid responses and expansion of CD8 T cells that is not maintained if the costimulatory molecule is removed may also suggest that more cells are initially recruited by islets expressing CD80 as previously suggested by Allison et al. (25). Diabetes in the NOD mouse is caused by a heterogeneous group of cells and the ability of CD80 to lower the threshold of activation of lower avidity cells could allow for recruitment of these lower avidity cells to the islet and in situ expansion of cells. However, the low-avidity cells, although stimulated to full activation in response to CD80 stimulation, are not able to maintain the level of activation important for cytotoxicity in the absence of costimulatory molecules. Thus, on encounter of low levels of Ag in the absence of the costimulatory molecule, such as in the case of transplantation with islets that do not express CD80, or in secondary transfer when the recipients do not express CD80, the low-avidity CD8 T cells are unable to further contribute to the disease process. Our results also suggest that these CD80-stimulated T cells are more susceptible to inhibition by CD4+CD25+ regulatory T cells. This may further imply that the islet destructive repertoire is different in the RIP-CD80 mice compared with the RIP-CD86 mice and nontransgenic NOD mice.

In conclusion, our studies have shown that when CD80 and CD86 stimulation are separated and tested in vivo, both CD80 and CD86 can stimulate activating and inhibitory responses but CD80 induces, overall, stronger responses. Thus, CD80 targeted locally may be useful in the context where CD80 remains expressed and continues to be active (39) to stimulate lower avidity T cells and overcome inhibition by CD4+CD25+ T cells. Although once activated, effector cells may not require continued costimulation under all circumstances, this depends on continued lower threshold of activation and is likely also to be dependent on the level of Ag expression. Overall, while CD86 does not stimulate an initial response as strongly as CD80, there is greater sustained activity that is seen, even in the absence of continued costimulation. These functions have implications for the engineered use of costimulatory molecules in altering immune responses in a therapeutic setting.

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Disclosures
The authors have no financial conflict of interest.

References


